FOUR CLINICAL VARIANTS OF CONGENITAL ADRENAL HYPERPLASIA

BY

W. HAMILTON and M. G. BRUSH

From the University Department of Child Health and the Royal Hospital for Sick Children, Glasgow, and the University Department of Steroid Biochemistry and the Royal Infirmary, Glasgow

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Three clinical types of congenital adrenogenital virilism due to adrenal hyperplasia have now been well defined. These are simple virilization, virilization with excessive sodium loss and danger to life and virilization combined with hypertension. Clinical subvariants have also been described in association with hypoglycaemia (White and Sutton, 1951; Wilkins, Crigler, Silverman, Gardner and Migeon, 1952), with periodic fever (Gonzales and Gardner, 1956; Gardner and Migeon, 1959) and with the late onset of sodium loss (Cara and Gardner, 1960). More recently Bongiovanni (1962) has described a variant that, when occurring in males, is characterized by incomplete masculinization, hypospadias and usually severe sodium loss, although in females it leads to the familiar virilism.

The treatment of congenital adrenal hyperplasia has been much simplified since Wilkins, Lewis, Klein and Rosenberg (1950) showed that cortisone decreased the urinary 17-oxosteroids. Deoxycorticosterone acetate (D.C.A.) in conjunction with cortisone and increased dietary salt usually control sodium loss. However, some cases have proved to be resistant to cortisone, though not to the newer steroid analogues, while others have failed to respond to seemingly adequate steroid replacement.

The adrenocortical enzyme defects underlying the principal types of adrenogenital virilism have been extensively investigated and the subject has been reviewed recently (Bongiovanni and Eberlein, 1961). However, the occurrence of further clinical subvariants indicates the need for separate clinical and biochemical elucidation of each case as a guide to adequate or optimum treatment. The following four cases serve to emphasize these facts. As far as is known, Case 4 is the first of its kind to be described in this country.

Case Reports

Case 1. This child, born January 27, 1954, was of ambiguous sex having a curved phallos with an opening at the tip. Both this and another opening on the perineum admitted a probe to a depth of 1 cm. The scrotum was bifold and the testes were not palpated.

When 5 weeks of age, a skin biopsy and buccal smear examined for sex chromatin indicated that the child was female. The urinary 17-oxosteroids were reported as 0.4 mg. per day. When 2 years of age the perineum was opened up in the midline and the urethral and vaginal orifices were found to open on to a small vestibule. The incision was sutured to leave the vestibule open and on healing the perineum looked remarkably normal for a female child.

During the following four years, the only feature noted was progressive clitoral enlargement. When the girl was 6½ years of age, the clitoris was amputated, and at laparotomy, normal ovaries, tubes and uterus were seen. Histological examination of material from the gonads showed only ovarian tissue with normal primordial follicles and some cystic follicles. The clitoris continued to enlarge. Pubic hair and breast development appeared.

She was admitted to the Royal Hospital for Sick Children, Glasgow, on November 12, 1962, for further investigation. She was then 8 years 10 months. Her height was 137 cm. (average 126 cm.) and she weighed 29.5 kg. (average 25 kg.). Apart from an enlarged clitoris and pubic hair (Fig. 1) she was a healthy girl. There was no history of vaginal bleeding. Bone radiography showed a normal bone age. A high urinary excretion of steroid metabolites (Table 1) was found, while the daily oestrogen levels were: oestrone 2.3 µg.; oestradiol 1.1 µg.; oestriol 1.9 µg. (These values are within normal range for age of patient.)

It was decided at this time to suppress adrenocortical function with prednisolone (Δ1-hydrocortisone) 10 mg. daily.

Summary. A 9-year-old female pseudohermaphrodite with adrenocortical hyperplasia was treated first by excision of the clitoris and then by replacement therapy. At present she is having prednisolone 5 mg. daily.
CONGENITAL ADRENAL HYPERPLASIA

Case 2. She was born on May 16, 1954, admitted to the Royal Hospital for Sick Children, Glasgow, at 2 weeks in acute adrenal failure. The genitalia are shown in Fig. 2. Diagnosis, immediate treatment and early management have previously been reported (Coleman and Arneil, 1958). Substitution therapy was with cortisone 25 mg. thrice daily, deoxycorticosterone acetate (D.C.A.) 200 mg. implants at 6-month intervals.

**TABLE 1**

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Age</th>
<th>Before Treatment</th>
<th>After Treatment</th>
<th>Drug and Dosage (mg./day)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>17-OS 17-OGS 17-OHCS 21-deoxyketols P'triol R</td>
<td>17-OS 17-OGS 17-OHCS 21-deoxyketols P'triol R</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>9 yrs.</td>
<td>6:9 5:6 10:3 4:7 30*</td>
<td></td>
<td>2:1 1:5 2:9 1:4</td>
</tr>
<tr>
<td>2</td>
<td>3 yrs. 7 yrs. 71 yrs.</td>
<td>4:0</td>
<td>48</td>
<td>2:5 5 DX 6 DX 4 DX 1 9a F 0:4</td>
</tr>
<tr>
<td>3</td>
<td>15 mths. 21 mths. 26 mths. 42 mths.</td>
<td>5:5 6:0 9:0 3:0 2</td>
<td>1:3 E 50 E 25 E 50 DX 2</td>
<td>2:4 4:7 8:4 3:1 1:0 0:12</td>
</tr>
<tr>
<td>4</td>
<td>Birth 2 days 3 days 6 days 7 days 20 days</td>
<td>4:8 4:3 2:9 5:2 3:6 2:1</td>
<td>0:048 0:028 0:049 E 20</td>
<td>1:8 8:4 11:3 2:9 1:8</td>
</tr>
</tbody>
</table>

P'triol, pregnanetriol; R, 11 deoxy/11-oxygenated-17-oxosteroids.
$\Delta^1$-F, prednisolone; DX, dexamethasone; E, cortisone; 9aF, 9α-fluorohydrocortisone.
* Overestimate due to presence of non-specific chromogens.
and extra dietary salt. By 2 years of age she had acquired a voracious appetite for salt and was taking 20-25 g. daily. This was associated with considerable thirst and polyuria, but further D.C.A. was not given. At 3 years of age, prednisolone 20 mg. daily was given to replace cortisone, since the urinary excretion of 17-oxosteroids was then 3-4 mg. per day. Salt addiction persisted, but there was no further virilization. When 4½ years old her bone age was equivalent to 1½ years and because of this, prednisolone was gradually reduced to 5 mg. daily. During the following year she had two adrenal crises, precipitated by mild respiratory infections. These were readily controlled and on each occasion D.C.A. implants of 100 mg. were made.

She was admitted when 7 years old (1962) for adrenal and gynaecological assessment. She was then 108 cm. (average 118·4 cm.) and weighed 20·5 kg. (average 22·1 kg.). To obtain baseline urinary levels, prednisolone was withdrawn but salt was allowed. Inadvertently, extra salt was omitted from the diet and within 48 hours a salt-losing crisis ensued. At this period urinary pregnanetriol was found to be at the high level of 48 mg. per day, using the method of Fotherby and Love (1960). To exclude the presence of non-specific chromogens estimating as pregnanetriol, the analysis was checked by the procedure of Cox (1959) and was found to be correct.

Dexamethasone (9-α-fluoro-16-α-methylprednisolone) 6 mg. daily and salt 20 g. daily were given with immediate good effect. Over 21 days dexamethasone was reduced to 1 mg. daily, but the high salt intake continued. It will be seen from Table 1 that urinary pregnanetriol excretion rapidly fell to normal levels with the commencement of suitable steroid therapy. She continued on this régime for two months and became extremely moon-faced and obese, although without hirsutism or hypertension.

In an attempt to control the polyuria and polydipsia induced by her large appetite for salt, it was decided to replace dexamethasone with 9-α-fluorohydrocortisone. The dosage of this drug was increased gradually from 0·1 mg. to 0·4 mg. daily, when the child abruptly stopped eating salt and took food without added salt. Polyuria and polydipsia ceased; her blood pressure remained about 100/70 mm. Hg. Pregnanetriol excretion was then 125 µg. per day (Table 1).

**Summary.** A 7-year-old female pseudohermaphrodite with adrenocortical hyperplasia accompanied by excessive sodium loss, was first treated with cortisone, D.C.A. and extra dietary salt and later with prednisolone and extra salt. At present she is well controlled on 9-α-fluorohydrocortisone 0·45 mg. daily without extra salt. She has grown 6 cm. in eight months and has lost the Cushing-type appearance.

**Case 3.** This male child was born on July 8, 1958, and was the fourth child of a full cousin marriage. The neonatal period was uneventful. He grew rapidly and when he was 1 year and 3 months his parents observed early pubic hair and penile enlargement (Fig. 3). Clinical examination at this time showed a child, tall for his age, muscular and with slight breast enlargement. Bone age was 4 years. Blood pressure was normal. Urinary excretion of 17-oxosteroids was 5·5 mg. per day. He was given cortisone 50 mg. daily for one month and thereafter 25 mg. daily. 17-oxosteroid output was then 1·3 mg. per day.

When seen six months later, growth had proceeded unchecked. His weight age was 4½ years, height age 2½ years and bone age 6 years. Urinary steroid excretion was again raised (Table 1). Cortisone was increased to 75 mg. daily. His parents alleged that this high dosage was maintained till he was next seen at 3 years of age. His bone age was then equivalent to 10 years. Under strict in-patient supervision cortisone was increased over six months to 300 mg. daily. Nevertheless, his bone age continued to advance to 11 years. Cortisone was then withdrawn abruptly without obvious ill effects. He was then given dexamethasone 2 mg. daily plus cortisone 50 mg. daily.

**Special Investigations.** Skull radiographs showed a normal pituitary fossa. Intravenous pyelogram and presacral gas insufflation of the retroperitoneum revealed kidneys and adrenal glands normal in size, shape and position. Blood corticotrophin was 5·71 milli units per 100 ml. whole blood (mean normal adult value 0·71 m.u. per 100 ml. blood). It will be seen from Fig. 4 that the adrenal glands were suppressed by corticosterone and dexamethasone. He was therefore maintained on dexamethasone 2 mg. daily and cortisone 50 mg. daily.

**Summary.** A male with pubertas praecox and advanced ossification due to adrenocortical hyperplasia. Cortisone in daily doses of 300 mg. did not suppress
adrenal activity. Control was obtained with dexamethasone 2 mg. daily to which cortisone 50 mg. daily was added to maintain adequate sodium retention.

Case 4. This male child was born on October 10, 1962, and was the third child of healthy unrelated parents.

![Image of child](image)

**Fig. 5.—Case 4, at birth. Genitalia suggest female virilism, but nuclear sex is male.**

The first child, a male, died in another hospital when 1 month old and at autopsy he was found to have congenital adrenal hyperplasia. The genitalia were reported to be normal.

At birth in the Royal Maternity Hospital, Glasgow, he was seen to be cryptorchid, with scrotal-labial folds. The urethra opened at the tip of a small hooded phallus (Fig. 5). The serum electrolytes in cord blood and subsequently in venous blood are shown in Table 2. A tentative diagnosis of female virilism was made but urinary pregnanetriol output during the first three days of life was 48, 28 and 49 µg. per day (these figures are probably normal for a newborn child, although the

**Table 2**

<table>
<thead>
<tr>
<th>Date</th>
<th>Drug (mg./day)</th>
<th>Na mEq per litre</th>
<th>K mEq per litre</th>
<th>Cl mEq per litre</th>
<th>Urea (mg./100 ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.10.62</td>
<td>D.C.A. 100</td>
<td>135</td>
<td>6</td>
<td>101</td>
<td>28</td>
</tr>
<tr>
<td>11.10.62</td>
<td>E: 15</td>
<td>143</td>
<td>6</td>
<td>103</td>
<td>30</td>
</tr>
<tr>
<td>15.10.62</td>
<td>E: 20</td>
<td>140</td>
<td>6</td>
<td>101</td>
<td>33</td>
</tr>
<tr>
<td>19.10.62</td>
<td>27 D.C.A.: 10</td>
<td>125</td>
<td>4</td>
<td>95</td>
<td>73</td>
</tr>
<tr>
<td>26.10.62</td>
<td>31 D.C.A.: 3</td>
<td>130</td>
<td>9</td>
<td>92</td>
<td>73</td>
</tr>
<tr>
<td>8.11.62</td>
<td>1.11.62 NaCl: 3</td>
<td>126</td>
<td>10</td>
<td>90</td>
<td>73</td>
</tr>
<tr>
<td>9.11.62</td>
<td>3.11.62 E: 25</td>
<td>122</td>
<td>8</td>
<td>102</td>
<td>70</td>
</tr>
<tr>
<td>10.11.62</td>
<td>5.11.62 D.C.A.:</td>
<td>126</td>
<td>5</td>
<td>102</td>
<td>73</td>
</tr>
<tr>
<td>16.11.62</td>
<td>19.11.62 D.C.A.:</td>
<td>134</td>
<td>6</td>
<td>118</td>
<td>90</td>
</tr>
<tr>
<td>29.11.62</td>
<td>4.12.62 E: 50</td>
<td>140</td>
<td>5</td>
<td>108</td>
<td>60</td>
</tr>
<tr>
<td>10.1.63</td>
<td>16.1.63 D.C.A.:</td>
<td>134</td>
<td>7</td>
<td>100</td>
<td>68</td>
</tr>
<tr>
<td>24.1.63</td>
<td>19.2.63 D.C.A.:</td>
<td>140</td>
<td>6</td>
<td>105</td>
<td>36</td>
</tr>
</tbody>
</table>

D.C.A., deoxycorticosterone acetate.

E, cortisone.

Dx, dexamethasone.

C.B., dexamethasone.

E, dexamethasone.

D.C.A., deoxy cortisol acetate.

![Graph](graph)

**Fig. 4.—The response of urinary 17-OHCS in Case 3 to orally administered corticosterone (100 mg. daily) and dexamethasone (1·5 mg. daily).**

![Graph](graph)

**Fig. 5.—Case 4, at birth. Genitalia suggest female virilism, but nuclear sex is male.**
improvement was only obtained when cortisone was increased to 100 mg. daily. After a prolonged period at this high level he was discharged from hospital when 14 weeks old. He was then receiving, daily, cortisone 50 mg., D.C.A. 1 mg. intramuscularly, and salt 3.5 g.

When seen as an out-patient, he had progressed well, and when aged 6 months he weighed 6.46 kg.; blood pressure was 100 mm. Hg. (flush technique).

Meanwhile chromatographic separation of steroids from pretreatment urine was in progress and the results are shown in Table 3. The principal finding was a high excretion of dehydroepiandrosterone (35 μg. per day; normal 0-7 μg. per day). This indicates at least a partial 3-β-hydroxysteroid (3-β-ol) dehydrogenase deficiency.

Summary. A male child, observed from birth till 6 months of age, with congenital hyperplasia of the adrenal cortex. Urinary steroid analyses indicated a partial block in the conversion of pregnenolone to progesterone due to a 3-β-ol-dehydrogenase deficiency. Clinically there were signs of adrenocortical insufficiency and improvement was achieved only by seemingly excessive doses of cortisone and D.C.A.

**Table 3**

<table>
<thead>
<tr>
<th>Steroid</th>
<th>μg./day</th>
</tr>
</thead>
<tbody>
<tr>
<td>Androsterone</td>
<td>25-7</td>
</tr>
<tr>
<td>Aetiocholanolone</td>
<td>91-9</td>
</tr>
<tr>
<td>Dehydroepiandrosterone</td>
<td>25-0</td>
</tr>
<tr>
<td>Tetrahydrocortisone</td>
<td>347-7</td>
</tr>
<tr>
<td>Pregnanetriol</td>
<td>42-0</td>
</tr>
<tr>
<td>Δ⁵-Pregnenediolone</td>
<td>76-2</td>
</tr>
</tbody>
</table>

Methods for Steroid Analysis

Since these investigations have been undertaken over the past 11 years, the chemical methods employed and the number of steroid groups estimated have varied. Urinary 17-oxidosteroids were determined at first according to Tomsett (1949) and later by a modification of the M.R.C. method (M.R.C. Committee on Clinical Endocrinology, 1951). For urinary 17-hydroxycorticosteroids, 17-oxygenic and 21-deoxysteroids the methods of Appleby, Gibson, Norymberski and Stubbs (1955) and Appleby and Norymberski (1955) were used. The ratio (R) of the 11-deoxy to the 11-oxygenated-17-oxosteroids derived from the 17-oxosteroids was found by the method of Morris (1959). This method estimates pregnanetriol in urine as a 17-oxygenic steroid, the oxidation of pregnanetriol to aetiocholanolone being accomplished by sodium bismuthate. In urines containing excess quantities of pregnanetriol, 11-oxo-aetiocholanolone will be formed after sodium bismuthate oxidation. A normal R value in congenital adrenal hyperplasia will only be significant if it is known that pregnanetriolone is not increased.

The Fotherby and Love (1960) procedure for the estimation of urinary pregnanetriol was used, although in certain instances paper chromatographic examination by the Cox (1959) technique showed the presence of some non-specific chromogenic material in the final extract. In such cases the Cox (1959) technique was used to give a specific semiquantitative estimation of pregnanetriol. Urinary oestrogens were estimated by the method of Brown (1955). Fractionation of individual urinary steroids in Case 4 was by the techniques described by Birchall, Cathro, Forsyth and Mitchell (1961).

**Discussion**

It is now recognized that congenital adrenal hyperplasia is a genetically determined enzymatic defect in the biosynthesis of hydrocortisone (cortisol). The common defect involves the C-21-hydroxylation of 17-α-hydroxyprogesterone. In the hypertensive form of congenital adrenal hyperplasia the defect involves the C-11-hydroxylation of 11-deoxycortisol (Compound S). Recently Bongiovanni (1962) has shown that in some cases the conversion of pregnenolone to progesterone is impaired (Fig. 6). This block, early in the pathway, is due to a deficiency of the enzyme 3-β-hydroxysteroid dehydrogenase (3-β-ol-dehydrogenase).

A partial C-21-hydroxylase defect results in the simple virilizing form of the disease, but when the defect is complete, sodium loss is present (Eberlein and Bongiovanni, 1960). The principal sodium-retaining corticosteroids, aldosterone and 11-deoxycorticosterone (DOC), carry a C-21-hydroxyl group. Thus the difference between the simple virilizing type and the sodium-losing type is quantitative rather than qualitative and only one of degree. Earlier, the same authors pointed out that in congenital adrenal hyperplasia due to C-11-hydroxylase deficiency, the hypertension was due to an abnormal production of DOC (Eberlein and Bongiovanni, 1955).

In the absence of 3-β-ol-dehydrogenase, production of aldosterone, DOC and cortisol is impossible and the condition is probably incompatible with life beyond a few days or weeks. A partial deficiency of the enzyme, however, results in reduced amounts of active corticosteroids, and five of the six cases so far described (Bongiovanni, 1962) have been 'salt-losers'. 3-β-ol-hydrogenase also plays a part in the gonadal synthesis of testosterone, and a deficiency of the enzyme during the embryonic period could result in incomplete masculinization of the male foetus (Bongiovanni, 1962). This is likely to be the explanation for the incomplete masculinization in Case 4.

The pattern of urinary steroid metabolites differs in each of these enzyme deficiencies. With C-21-hydroxylase deficiency, pregnanetriol and 11-oxypregnatriol (pregnanetriolone) are increased, while
with the C-11-hydroxylase deficiency the metabolites of compound S and DOC (i.e. tetrahydro-S and tetrahydro-DOC) and pregnanetriol are found but not pregnanetriolone. In 3-β-ol-dehydrogenase deficiency, metabolites of pregnenolone appear in the urine as well as other substances with the 3-β-hydroxysteroid (Δ5 3-β-ol) structure. Included in this group is dehydroepiandrosterone (DHEA), a precursor of testosterone.

Case 1 is an example of partial C-21-hydroxylase deficiency showing virilism as the only clinical feature. Although the clitoris was amputated its growth continued until effective adrenal suppression was achieved with prednisolone.

Case 2 demonstrates the result of a complete C-21-hydroxylase deficiency, namely virilism and salt loss. Originally she required D.C.A. in addition to cortisone, but subsequently did well on prednisolone alone while permitted to take salt freely. At the time of writing she is adequately controlled on 9-α-fluorohydrocortisone 0-45 mg. daily and has no further desire for excess salt. The daily excretion of pregnanetriol is also normally low.

Compared with cortisol, 9-α-fluorohydrocortisone is more active as a glucocorticoid by 10 times and as a mineralocorticoid by 800 times. It thus appears that control of sodium loss is a more important factor in adrenal suppression than replacement of glucocorticoids.

In Case 3, the adrenal overactivity was not suppressed by high doses of cortisone. This was evidenced chemically by a continued high urinary excretion of steroid metabolites and clinically by advancing ossification and growth. Green, Cleveland and Wilkins (1961) have observed escape from suppression with cortisone and have attributed this to the transient action of oral cortisone. They report the successful use of triamcinolone (9-α-fluoro-16-α-hydroxy prednisolone) when cortisone had failed. Cox (1962) found that pregnanetriolone disappeared from the urine in congenital adrenal hyperplasia during treatment with cortisone more slowly than pregnanetriol. A normal R value (Morris, 1959; Hill, 1960) found in congenital adrenal hyperplasia during treatment need not therefore indicate good control. In our case
cortisone was given four times daily to reduce the tendency to escape, but the urinary pregnanetriol and pregnanetriolone remained comparatively high. Absorption from the gut seemed adequate as indicated by the high urinary 17-hydroxycorticosteroids during treatment. Good adrenal suppression was achieved with dexamethasone, but since this substance has no mineralocorticoid activity, a small amount of cortisone was included in maintenance therapy to assist in sodium retention.

Case 4 is an example of the salt-losing type of congenital adrenal hyperplasia in a male with cryptorchidism. He was resistant to replacement therapy in ‘physiological’ doses, but responded to a much increased dosage of cortisone and D.C.A. Bongiovanni (1961, 1962) was the first to report six cases of this new syndrome, five of which were ‘salt-losers’, and these five died within the first three months of life. Three of the six cases were male with hypospadias, and he suggests that the failure to complete masculine development is in keeping with a deficiency of 3β-ol-dehydrogenase. The enzyme is known to have a part in the gonadal biosynthesis of testosterone. The partial nature of the enzyme deficiency in our case is clear since significant quantities of tetrahydrocortisone and normal amounts of pregnanetriol were present in the urine (Table 3).

Summary

The clinical types of adrenogenital virilism are reviewed.

Four such cases, due to congenital adrenal hyperplasia, are described in detail. The biochemical findings and treatment are outlined.

In discussion, the enzyme deficiencies in the biosynthesis of hydrocortisone, now recognized as the cause of the condition, are considered, and the resultant patterns of urinary steroid metabolites are indicated.

The report includes an example, believed to be the first described in this country, of congenital adrenal hyperplasia due to a deficiency of 3β-hydroxysteroid dehydrogenase.

We are grateful to Professor J. H. Hutchison for permission to publish these cases and for his constant help and encouragement. Thanks are also due to Dr. Beryl Davis for estimating blood corticotrophin, Dr. Eileen E. Hill for monitoring the early R values in Case 3, Dr. J. B. Brown for the oestrogen levels in Case 1, and Dr. F. L. Mitchell for chromatographic separation of individual urinary steroids in Case 4. Part of the expense of these investigations was defrayed by the Rankin Funds of the University of Glasgow.

REFERENCES


